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Kim, Yejin

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# **Anti-cancer Drug Synergy Prediction in Understudied Tissues using Transfer Learning**

Yejin Kim<sup>1</sup>, Shuyu Zheng<sup>2</sup>, Jing Tang<sup>2</sup>, W. Jim Zheng<sup>1</sup>, Zhao Li<sup>1</sup>, Xiaoqian Jiang<sup>1</sup>

<sup>1</sup>School of Biomedical Informatics, University of Texas Health Science Center at Houston, Houston, Texas, United States

<sup>2</sup>Research Program in Systems Oncology, Faculty of Medicine, University of Helsinki, Helsinki, Finland

Correspondence should be addressed to Y.K. ([yejin.kim@uth.tmc.edu](mailto:yejin.kim@uth.tmc.edu); 7000 Fannin Street, Houston, Texas; 713.500.3998)

## ABSTRACT

**Motivation:** Drug combination screening has advantages in identifying cancer treatment options with higher efficacy without degradation in terms of safety. A key challenge is that the accumulated number of observations in in-vitro drug responses varies greatly among different cancer types, where some tissues are understudied than the others. Thus, we aim to develop a drug synergy prediction model for understudied tissues as overcoming data scarcity problems.

**Methods:** We collected a comprehensive set of genetic, molecular, phenotypic features for cancer cell lines. We developed a drug synergy prediction model based on multi-task deep neural networks to integrate multi-modal input and multiple output. We also utilized transfer learning from data-rich tissues to data-poor tissues.

**Results:** We showed improved accuracy in predicting synergy in both data-rich tissues and understudied tissues. In data-rich tissue, the prediction model accuracy was 0.9577 AUROC for binarized classification task and 174.3 mean squared error for regression task. We observed that an adequate transfer learning strategy significantly increases accuracy in the understudied tissues.

**Conclusions:** Our synergy prediction model can be used to rank synergistic drug combinations in understudied tissues and thus help to prioritize future in-vitro experiments. Code is available at <https://github.com/yejinjkim/synergy-transfer>.

## INTRODUCTION

Discovering synergistic drug combination in cancer treatment has been widely studied as it identifies candidates with higher efficacy without degradation in terms of safety.<sup>1-4</sup> Many previous studies show great promises in finding potent drug combinations that cannot be identified by one drug - one target approaches.<sup>5-10</sup> A considerable amount of drug synergy prediction methods has been investigated including pharmacokinetic and pharmacodynamic methods,<sup>10,11</sup> post-treatment transcriptome,<sup>12,13</sup> feature-based machine learning,<sup>5,7,8,14,15</sup> neural networks with multi-modal data sources,<sup>4,16</sup> and large community efforts.<sup>17,18</sup> These studies utilize various features such as molecular profiles, chemical structure, pre- or post-treatment transcriptome. Their prediction target usually remains in a specific pathway, cell lines, or tissue. These studies are usually based on single databases and the prediction was also made within the database.<sup>3-5,18</sup> A systematic integration of multiple drug synergy databases has been recently proposed,<sup>19,20</sup> which shed the light on developing unbiased drug synergy prediction models.

On the developing unbiased and generalizable drug synergy prediction model, a key challenge is an imbalance of synergy experimental data, in which the accumulated number of observations in in-vitro drug responses varies greatly for different tissues. Traditional methods target studying commonly observed tissues such as breast, kidney, skin, lungs.<sup>4,7,17,21-25</sup> These methods take known drug response data at certain cell lines and attempt to find other drugs responses at other cell lines within the same data-rich tissues. The understudied tissues include bone, prostate, and pancreas (Fig. 1a and 1b). A number of obstacles impede the drug response study in these tissues. For example, bone cancer drugs response has been poorly understood due to physical difficulty of culturing bone tissues as cell lines, the rarity of the tumors in sarcoma type, the difficulties of obtaining tumor tissue fragments from human patients for bone metastasis, and thus the limited number of cell models.<sup>26</sup> The lack of cell line models makes the high-throughput screening difficult, which in turn makes these tissues even more understudied. There is,

therefore, a critical need to develop a computational drug combination prediction tool for the understudied tissues.

Our goal is to develop a generalizable in-silico drug synergy prediction model in understudied tissues. The most critical challenge is data scarcity. Understudied tissues inevitably lack enough training data; they have a limited number of experimental observations and also lack important features such as post-treatment transcriptome. A potential way to mitigate this data scarcity problem is to utilize information from the data-rich tissues to the data-poor tissues as these different tissues share biological commonality partly in terms of gene expression and therefore respond drugs in similar ways.<sup>27</sup> Several previous studies support that anti-cancer drug sensitivity in cell lines is not tissue-specific and that tissue-specific drugs can bring additional benefits to other tissues.<sup>28,29</sup> Accordingly we proposed to utilize interaction between drugs and cell lines learned from the data-rich tissues to help increase the performance of the data-poor tissue using *transfer learning*. Transfer learning is to transfer knowledge (in the form of parameters in machine learning model) from previously learned model (teacher model) with large data to new model (student model) with fewer data. Several methods have been proposed to utilize transfer learning in drug sensitivity prediction to link complementary data sources, different modality of auxiliary feature, and different disease.<sup>30,31</sup> To learn generalizable and thus transferable knowledge on drug target and cell line gene expressions, we integrated all types of available tissues in one model with a large dataset (i.e., 4,150 drugs; 112 cell lines; 710,242 monotherapy sensitivity; 1,175,220 combinations synergy from 15 different synergy studies). We maximally utilized various types of multi-modal inputs (molecular, genetic, phenotypic features) and multiple outputs (drug sensitivity and synergy) using flexible neural networks models, and we validated our ranking models and drug combinations with independent external databases.

## **MATERIALS**

## Drug sensitivity and synergy

We used a publicly available large-scale drug synergy database from DrugComb Portal,<sup>19</sup> which combines 15 drug synergy studies into one integrated database including O’Neil, Cloud, ALMANAC, NCATS Matrix, Forcina, Mathews, Phelan, Wilson, and Yohe (Table S1).<sup>3,32–38</sup> The number of unique drugs and cell lines were 4,150 and 112, respectively (Fig. 1). There are 2,843 experimental drugs and 1,307 FDA-approved drugs. There were a total of 710,242 monotherapy sensitivity (a pair of drug and cell line) and 466,259 drug combinations synergy (a triplet of drug1, drug2, and cell line). For monotherapy sensitivity we calculated relative inhibition (RI) from dosage combination matrix (Supplementary 1). For drug combinations synergy we used Loewe synergy score (Fig. 1), which ranged from -100 (antagonistic effect) to 75 (strong synergistic effect).<sup>39</sup> Loewe synergy score is to quantify the excess over the expected response if the two drugs are the same compound.<sup>40,41</sup> We selected Loewe score for comparison with the baseline study.<sup>4</sup>

## Drug’s features

We extracted the drug's molecular and genomic features. For molecular features, we used both Molecular ACCess System (MACCS) fingerprints<sup>42</sup> and native chemical compounds. MACCS fingerprints contain 166 chemical structures such as the number of oxygen, S-S bonds, and rings. We used RDKit (<http://www.rdkit.org>) to extract MACCS fingerprints. In addition, we represented drugs as a native chemical structure using SMILES. SMILES is a linear notation to represent chemical compound uniquely; in the SMILES representation atoms are represented as their atomic symbols (e.g., c for carbon); special characters are also used to represent relationship (e.g., “=”: double bonds; “#”:triple bonds; “.”:ionic bond; “:”: aromatic bond)<sup>43</sup>. SMILES can provide richer features space that strictly represent functional substructures and express structural differences such as compound’s chirality<sup>44</sup>. We used the state-of-the-art natural language processing model, Transformer, to encode the SMILES

sequence.<sup>45</sup> For genomic features, we integrated three different drug databases — DrugBank,<sup>46</sup> Therapeutic Targets Database (TTD),<sup>47</sup> and NIH-LINC<sup>48</sup> — for complete view of drug’s target genes (Table S2). We filtered non-human target genes in TTD.

### **Cell line’s features**

We used the cell line’s cancer type and genomic features. There were 14 tissues including lung, ovary, and skin and 14 cancer types including carcinoma, adenocarcinoma, and melanoma (Table S3). We also extracted gene expression profiles by Fragments Per Kilobase of transcript per Million reads mapped (FPKM) from CCLE and COSMIC.<sup>49,50</sup> We found 75 cell lines with 37,260 genes from Broad Institute and 25 cell lines with 35,004 genes from Sangar (Table S2). In total, we found 88 cell lines with 22,586 genes after excluding zero-variance genes. We normalized the FPKM values into z-score in a gene-wise manner as FPKM varies greatly depending on gene. We only used baseline (before-treatment) gene expression profiles without after-treatment gene profiles because our objective is to test our models in understudied tissues, which rarely has after-treatment gene profiles.

## **METHODS**

### **Method overview**

Our transfer learning approach to enhance synergy prediction in data-poor tissue is to transfer parameters from a pre-trained general prediction model to the specific tissue’s prediction models. We first learned the general prediction model with data-abundant tissues (including brain, breast, colon, endometrium, hematopoietic and lymphoid, kidney, lung, ovary, and skin) from the pooled drug synergy databases containing the 15 different databases. We used neural networks as the prediction model due to its



flexibility in parameter sharing. The prediction model for the data-poor tissue was initialized with the pre-trained general model’s parameters and re-trained with its own data accordingly.

## General prediction model

Our objective is to predict whether unobserved drug combinations have synergistic effects in a given cell line and provide a list of combinations that researchers can prioritize for experiments. We formulated it as a prediction problem. Given an experimental block  $\mathcal{B}_c = \{\mathcal{D}_1, \mathcal{D}_2, \mathcal{D}_3\}$  of drug combination  $(\mathcal{D}_i, \mathcal{D}_j)$  and cell line  $\mathcal{C}_c$ , the prediction model is a function  $\mathcal{F}$  such that

$$\mathcal{F}_{\mathcal{B}_c} \approx \mathcal{F}(\mathcal{D}_i, \mathcal{D}_j, \mathcal{C}_c),$$

where  $\mathcal{F}_{\mathcal{B}_c}$  is the synergy score of the drug combination. In addition to the synergistic effect, we used the drug’s monotherapy sensitivity as another prediction objective to boost the synergy prediction. For monotherapy sensitivity in the cell line, the prediction model is  $\mathcal{G}$ :

$$\mathcal{G}_{\mathcal{D}_i} \approx \mathcal{G}(\mathcal{D}_i, \mathcal{C}_c) \text{ and } \mathcal{G}_{\mathcal{D}_j} \approx \mathcal{G}(\mathcal{D}_j, \mathcal{C}_c),$$

where  $\mathcal{G}_{\mathcal{D}_i}$  is the sensitivity score of drug  $\mathcal{D}_i$  in cell line  $\mathcal{C}_c$ . We jointly optimize sensitivity and synergy as multi-task learning. To incorporate multimodal sources of inputs and formulate nonlinear relationships between chemical and genomic features, we used deep neural networks as the prediction function. Our prediction model consists of drug encoders (Fig. 2a), cell line encoder (Fig. 2b), and merging layers (Fig. 2c) for final prediction in an end-to-end manner. Using the estimated drug response, we can rank drug combinations at a cell line that are expected to have synergistic effect.

## Drug encoder

Drug encoder learns an embedding representation of each drug  $\mathcal{D}_i$  or  $\mathcal{D}_j$ . Each drug’s features are {MACCS fingerprints, canonical SMILES, target genes} (Table 1). Binary indicators of MACCS

fingerprints were used as raw input. Binary vectors of target genes were represented as compressed embedding with a single layer of denoising autoencoder.<sup>51</sup> As SMILES has variable length, we need more careful consideration. We used Transformer encoder, a natural language processing model that convert the sequence into a representation.<sup>45</sup> Each symbol in SMILES were first represented as one-hot vectors with size of (#SMILES length \* #unique symbols), where #unique symbols was 48 and the maximum SMILES length was 288. The symbol's one-hot vector was then represented as embedding. This embedding sequence was fed into a separate Transformer encoder, which consists of a multi-head attention layer and feed-forward layer with repeated residual connections. Once we derived all the embedding representations, they were concatenated into one and fed into two feed forward layers with Relu activation and dropout (Fig. 2a).

**Table 1.** Multimodal input and multi-task output to predict drug response. SMILES=Simplified molecular-input line-entry system. FPKM = Fragments Per Kilobase of transcript per Million reads mapped.

	Type	Size	Format
<b>Input</b>			
Drug features	MACCS fingerprint	166	Binary indicator vector
	Canonical SMILES	Variable length (max 288)	Sequences of characters (molecules and relation)
	Target gene	24,342 (Total number of unique genes)	Binary indicator vector
Cell line features	Cancer type	14	One-hot indicator vector
	Baseline gene expression	24,342 (Total number of unique genes)	Z-score of FPKM
<b>Output</b>			

Monotherapy sensitivity	Relative inhibition		Continuous value or binarized value at 50
Combination synergy	Loewe synergy score		Continuous value or binarized value at 30

### Cell line encoder

Cell line encoder learns an embedding representation of each cell line  $\square_{\square}$ . Each cell line's features are {cancer type, gene expressions} (Table 1). To combine the multimodal inputs, we derived embedding from each source and merged into one embedding. The cancer types were represented as the same sized embedding. The gene expression of each cell line was represented as compressed embedding from the normalized FPKM values with 1 layer of denoising autoencoder. We concatenated all three embeddings into one and fed them into two feed forward layers with Relu activation and dropout (Fig. 2b).

Note that we did not include drug ID or cell line ID as an input. The IDs are useful in collaborative filtering techniques, which assumes that two drugs will react to a cell line similarly if these drugs have responded similarly to certain cell lines. In contrast, our focus was to derive transferable general knowledge in prediction models from data-abundant domain to data-scarce one. The cell lines between the two domains are disjoint to each other in most cases (cold start for all cases), and the data-scarce tissues usually do not have a past drug response history.

### Merging layers

For synergy prediction, we merged embedding representations of  $\square_{\square}$ ,  $\square_{\square}$ ,  $\square_{\square}$  into one and fed them into two feed forward layers with Relu activation. For sensitivity prediction, we merged embedding representation from  $(\square_{\square}, \square_{\square})$  and  $(\square_{\square}, \square_{\square})$ , respectively, and fed them into two feed forward layers (Fig. 2c).

## Training loss

We trained the model as multi-task learning that predicts synergy and sensitivity simultaneously. For combination synergy prediction, the training loss was mean squared error (MSE):

$$\mathcal{L}_{\text{synergy}} = || \hat{s}_{\text{synergy}} - \sigma(\mathbf{w}_{\text{synergy}} \cdot \mathbf{x}_{\text{synergy}}) ||^2$$

where  $\hat{s}_{\text{synergy}}$  is the synergy score. The training loss for monotherapy  $(\mathbf{w}_{\text{synergy}}, \mathbf{x}_{\text{synergy}})$  and  $(\mathbf{w}_{\text{sensitivity}}, \mathbf{x}_{\text{sensitivity}})$  was similarly defined:

$$\mathcal{L}_{\text{total}} = || \hat{s}_{\text{synergy}} - \sigma(\mathbf{w}_{\text{synergy}} \cdot \mathbf{x}_{\text{synergy}}) ||^2 + || \hat{s}_{\text{sensitivity}} - \sigma(\mathbf{w}_{\text{sensitivity}} \cdot \mathbf{x}_{\text{sensitivity}}) ||^2$$

for all drugs and cell lines in the training set. In addition to these regressions, we also developed

classification models. We first binarized the drug responses by setting a threshold. That is,  $\hat{y}_{\text{synergy}} = 1$  if

$\hat{s}_{\text{synergy}} > \tau_{\text{synergy}}$ ; 0 otherwise, and  $\hat{y}_{\text{sensitivity}} = 1$  if  $\hat{s}_{\text{sensitivity}} > \tau_{\text{sensitivity}}$ ; 0 otherwise. The

synergy and sensitivity threshold were set at 30 and 50 following the previous study.<sup>4,52</sup> Then the

classification model's training loss was binary cross entropy:

$$\mathcal{L}_{\text{synergy}} = -\hat{y}_{\text{synergy}} \log(\sigma(\mathbf{w}_{\text{synergy}} \cdot \mathbf{x}_{\text{synergy}})) - (1 - \hat{y}_{\text{synergy}}) \log[1 - \sigma(\mathbf{w}_{\text{synergy}} \cdot \mathbf{x}_{\text{synergy}})]$$

$$\mathcal{L}_{\text{sensitivity}} = -\hat{y}_{\text{sensitivity}} \log(\sigma(\mathbf{w}_{\text{sensitivity}} \cdot \mathbf{x}_{\text{sensitivity}})) - (1 - \hat{y}_{\text{sensitivity}}) \log[1 - \sigma(\mathbf{w}_{\text{sensitivity}} \cdot \mathbf{x}_{\text{sensitivity}})] - \hat{y}_{\text{synergy}}$$

$$\log(\sigma(\mathbf{w}_{\text{synergy}} \cdot \mathbf{x}_{\text{synergy}})) - (1 - \hat{y}_{\text{synergy}}) \log[1 - \sigma(\mathbf{w}_{\text{synergy}} \cdot \mathbf{x}_{\text{synergy}})]$$

where  $\sigma$  is a sigmoid function. We alternated the two optimization tasks with respect to synergy or sensitivity in every epoch. That is, we first minimized  $\mathcal{L}_{\text{synergy}}$  with all training batches and then switched to  $\mathcal{L}_{\text{sensitivity}}$  for the next training round. Optimizer was adaptive gradient descent with Autograd in PyTorch 1.3.0.

### **Fine-tuning to data-poor tissues**

We focused on transfer learning from the pooled data-rich tissues to bone and prostate (Fig. 3a), respectively. For the transfer learning experiments, we chose the bone and prostate tissue (number of observations > 1,000) over soft tissue, endometrium, and pancreas (number of observations < 1,000) (Fig. 1a), because they have too few observations to build their own prediction model for comparative experiments. We first pre-trained the general drug response prediction model with the data-rich tissues to learn the underlying mechanism between drugs and cell lines, and built a separate prediction model for specific tissue of interest (i.e., bone or prostate), transferred the pre-trained model parameters from the general model, and fine-tuned the specific tissue models' parameters. Possible fine-tuning strategies include i) retraining all the parameters, ii) fixing drug encoders & retraining cell line encoders and merging layers, iii) fixing drugs and cell line encoders & retraining merging layers, and iv) no retraining at all. We compared the accuracy of the four different strategies in each tissue and reported the best one.

### **Evaluation**

**Training and test set.** We evaluated the general and specific prediction models with cross and external validation. For cross validation, we randomly separated the pooled databases into train (80%) and test set (20%) (Fig. 3b). For external validation, we set aside one independent database as a test set and used the remaining pooled databases as a training set (Fig. 3b); the independent test set had only a few overlapping cell lines or drugs with that of the training set (e.g., nine cell lines and 36 drugs were common in training and test sets in data-rich tissues). Each data-poor tissue (bone and prostate) had four distinct cell lines, respectively (Fig 1ab); different tissues sometimes shared drugs, such as the bone tissue sharing 1,437 drugs and the prostate tissue sharing 127 drugs with the data-rich tissues.

**Accuracy measures.** We measured accuracy of the regression and classification task. The regression accuracy measure was the mean squared error (MSE); the classification accuracy measures were area under the receiver operating curve (AUROC) and area under precision-recall curve (AUPRC).

**Baselines.** We compared our accuracy with two baseline methods — i) DeepSynergy that utilizes deep neural networks using drug’s descriptors and cell line’s gene expression on O’Neil database<sup>4</sup> and ALMANAC+XGBoost that utilizes gradient boosting trees.<sup>5</sup> We also implemented the collaborative filtering method that uses drugs and cell lines ID as input to compare our contents-based approach with collaborative approach. Benchmark accuracy values of various machine learning methods (including support vector machines, gradient boosting machines, and random forest) are listed in the referred paper.<sup>4</sup> Note that each training or test set was a triplet  $\{\square_{\square}, \square_{\square}, \square_{\square}\}$  of the drug pair and cell line in corresponding tissue. We did not report sensitivity prediction accuracy because the train/test split was disjoint in terms of synergy triplets but sometimes overlapped in terms of sensitivity pairs. For example, let us say we split the data as  $\{\square_1, \square_2, \square_1\}$  in the training set and  $\{\square_1, \square_3, \square_1\}$  in the test set for synergy prediction. This split implies  $\{\square_1, \square_1\}, \{\square_2, \square_1\}$  in training and  $\{\square_1, \square_1\}, \{\square_3, \square_1\}$  in the test set for sensitivity prediction. Due to the overlapped pairs  $\{\square_1, \square_1\}$ , the sensitivity prediction accuracy can be too optimistic.

## RESULTS

### Prediction accuracy of general model

We evaluated the accuracy of our synergy prediction models. First experiment was to evaluate the accuracy of general models trained and tested with data-rich tissues (Table 2). We achieved  $\{0.9577$  AUROC,  $0.8335$  AUPRC,  $132.4$  MSE $\}$  for cross validation (randomly split from 15 pooled databases) and  $\{0.6376$  AUROC,  $0.5564$  AUPRC,  $742.1$  MSE $\}$  for external validation (i.e., trained with 14 pooled

databases; tested with O’Neil). The collaborative filtering method based on drugs and cell IDs showed lower accuracy compared to our multimodal-contents-based model. The lower accuracy in external validation compared to cross validation might be partly due to different experimental settings on measuring the drug responses scores and gene expression values across various synergy databases.

We trained and tested our model with O’Neil database in particular to compare our model’s accuracy with DeepSynergy.<sup>4</sup> Our model achieved significantly higher AUROC/AUPRC and lower MSE — DeepSynergy showed {0.90 AUROC, 0.59 AUPRC, 255.5 MSE}; whereas our proposed model showed {0.95 AUROC, 0.94 AUPRC, 181.7 MSE}. In addition, we also trained our model with the ALMANAC database to compare our accuracy with that of NCI-ALMANAC + XGBoost model.<sup>5</sup> Our model achieved {0.97 AUROC, 0.84 AUPRC, 75.4 MSE}; whereas the baseline XGBoost model achieved 77.8 MSE. The increased accuracy was mainly due to more diverse multi-modal features (such as target genes and native SMILES) and multi-task learning with the monotherapy sensitivity.

**Table 2.** Drug synergy prediction accuracy for data-rich tissues. Cross validation with randomly split train/test from the 15 pooled databases. External validation with the O’Neil for test and the remaining 14 pooled databases for training. For classification tasks, AUROC and AUPRC were computed after binarizing synergy at threshold 30. AUROC=area under the receiver operating curve; AUPRC= area under the precision-recall curve; MSE=mean squared error.

Train	Test	Methods	AUROC	AUPRC	MSE
15 pooled databases	15 pooled databases	collaborative filtering	0.9336	0.7484	151.9
14 pooled databases	O’Neil		0.6030	0.5081	706.0
O’Neil	O’Neil	Proposed method	0.9477	0.9388	181.7
ALMANA C	ALMANA C		0.9688	0.8376	75.4
15 pooled databases	15 pooled databases		0.9577	0.8335	132.4
14 pooled databases	O’Neil		0.6376	0.5564	742.1



### Prediction accuracy of specific model with transfer learning

In next experiments we moved our focus to understudied tissues. These experiments compared the four different strategies on transferring model parameters from data-rich tissues to bone and prostate respectively. We also compared accuracy with and without the transferred model parameters. We found that an adequate transfer learning increases accuracy in both bone and prostate (Table 3). In external validation of bone cell lines, we achieved {0.6647 AUROC, 0.4150 AUPRC, 297.4 MSE} without transfer learning and {0.8015 AUROC, 0.4676 AUPRC, 195.9 MSE} with transfer learning (transferred all parameters and no fine-tuned). In the external validation of prostate cell lines, we achieved {0.6505 AUROC, 0.5220 APPRC, 528.5 MSE} without transfer learning and {0.8542 AUROC, 0.7804 AUPRC, 327.9 MSE} with transfer learning (fixed drug and cell line encoders and retrained last merging layers).

**Table 3.** Drug synergy prediction accuracy for bone and prostate cancer (data-poor tissue) with different transfer learning strategies in cross or external validation.

Tissue	Train	Test	Transfer	AUROC	AUPRC	MSE
Bone	DIPG, ES (FAKi/AURKi), ES (Naampt_PARP)	DIPG, ES (FAKi/AURKi), ES (Naampt_PARP)	No transfer	0.8283	0.5922	61.3
			Transfer and retrain all parameters	0.8357	0.5904	48.4
	DIPG, ES (FAKi/AURKi)	ES (Naampt_PARP)	No transfer	0.6647	0.4150	297.4
			Transfer and no retrain	0.8015	0.4676	195.9
Prostate	ALMANAC, ONEIL	ALMANAC, ONEIL	No transfer	0.9775	0.8575	78.5

			Transfer and retrain all parameters	0.9928	0.9628	47.4
			No transfer	0.6505	0.5220	528.5
	ALMANAC	ONEIL	Fix encoders and retrain last merging layers	0.8542	0.7804	327.9

## Ranking

We listed predicted drug combinations for bone and prostate using the externally validated model. We selected top 20 combinations based on estimated probability of being synergistic (Table 4, S3). For the bone cell lines, among the top 20 five combinations were actually synergistic (5/20=25% hit) in the independent validation set (ES Naampt\_PARP). Considering the fact that only 1.56% of combinations are synergistic in bone, our model effectively ranked synergistic drug combinations. For the prostate cell lines, all top 20 combinations were actually synergistic in the independent O’Neil database (only 2.15% of combinations are synergistic in the prostate).

**Table 4.** Top 5 drug combinations in bone and prostate. The bone model was trained with DIPG and ES (FAKi/AURKi), and tested with ES (Naampt\_PARP). The prostate model was trained with ALMANAC and tested with O’Neil. “Hit” was marked if the predicted drug combinations show Loewe synergy score >30. Full list of top 20 drug combinations in Table S4.

Drug1	Drug2	Cell line	Hit?
Bone			

AZD1775	AZACITIDINE	TC-71	
ZINC34894448	ZINC34894448	TC-71	✓
ZINC34894448	ZINC34894448	EW-8	
AZD1775	AZACITIDINE	EW-8	
thapsigargin	thapsigargin	TC-71	✓
Prostate			
TOPOTECAN	BEZ-235	LNCAP	✓
GELDANAMYCIN	BEZ-235	LNCAP	✓
DOXORUBICIN	BEZ-235	LNCAP	✓
BORTEZOMIB	BEZ-235	LNCAP	✓
5-FU	BEZ-235	LNCAP	✓

## DISCUSSION

The objective of this study was to develop the drug combination synergy prediction model, which can be used even in the understudied tissue with less observation. To meet this end, we i) collected a comprehensive set of multimodal features from multiple databases, ii) integrated the multimodal features and multiple tasks using deep neural networks, iii) transferred the prediction model from data-rich tissues to data-poor one. As a result, the proposed model predicted drug synergy accurately with 0.8015 AUROC for bone and 0.8542 AUROC for prostate in external validation with an adequate transfer learning strategy.

Main contribution of our study is that we tackle understudied but critical tissues for drug synergy prediction. The difficulty of obtaining cell lines in these understudied tissues has limited the in-vitro experiments, which consequently become an obstacle to drug development in these tissues. As we focused on understudied tissues, we should avoid using post-treatment gene expression profiles as an input feature. The post-treatment gene expression is the most powerful feature to estimate drug response, but it is only accessible after drug compounds are tested in cell line.<sup>7</sup> Our drug response prediction model was able to successfully bypass the post-treatment features while achieving competitive accuracy.

Although our study's focus is on predicting synergy in understudied tissues, our model achieved improved accuracy even on general data-rich tissues than that of the baseline model. This increased accuracy is possibly due to the large-scale and multi-modal data with multi-task optimization. We used the 15 complementary drug synergy databases including the NCI-ALMANAC, Cloud, O'Neil, and NCATS Matrix. We incorporated diverse and comprehensive multi-modal sources of features. This large-scale data collection allowed us to maximize the power of data-driven computational models based on deep neural networks.

Our model was designed to predict combination synergy together with monotherapy sensitivity simultaneously. Here monotherapy sensitivity was an auxiliary output to boost accuracy of synergy prediction. A previous study uses monotherapy sensitivity as an input feature to predict synergy,<sup>7</sup> but understudied tissues do not have enough experimental observation including monotherapy response features. To utilize this partial input feature in data-rich tissues (not in data-poor tissues), we used the monotherapy sensitivity as an auxiliary output in data-rich tissue models so that we can still transfer the information to data-poor tissues via the model parameters when minor tissues do not have monotherapy sensitivity.

The major limitation of this study is that the general prediction model was not able to achieve high accuracy in external validation. A possible explanation for this might be that the various synergy studies and various tissues may have different experimental settings on measuring the drug responses and gene expressions values, thus validation on the mixed tissues in the independent synergy experiment could cause the discrepancy in the predicted and observed values.

In conclusion, our model is an end-to-end drug synergy prediction model that learns interaction between drugs and cell lines. Based on the fact that different tissues share common gene expression and therefore respond to drugs in similar ways, we used transfer learning from data-rich tissues to data-poor tissues to make the synergy prediction model work in data-poor tissues. As a future work, our drug prediction model for understudied tissues can potentially shed light onto other diseases that share drug targets and underlying mechanisms and offer a novel way of efficient and low-cost drug discovery.

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### **Competing Interests Statements**

No competing interests to declare.

### **Contributionship Statement**

X. J., W. Z. and J. T. provided motivation of this study;

S. Z., Z. L. prepared necessary databases;

J. T., S. Z., and W. Z. provided important insight for model development;

X. J. and Y. K. designed prediction model;

Y. K. implemented models and prepared manuscript;

S. Z., J. T., W. Z., and X. J. provided proof reading.

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## FIGURE LEGENDS

**Figure 1.** Summary statistics of the DrugComb database. (a) Number of drug combinations and (b) cell lines in each tissue. ‘Hem&lymp’ = hematopoietic and lymphoid tissue. Data-rich tissues include hematopoietic and lymphoid, lung, skin, ovary, kidney, brain, breast, and colon, which have more than 20,000 drug combination tests with more than 5 different cell lines per each tissue. Data-poor tissues include prostate and bone, which have less than 15,000 drug combination tests within 3 or 4 cell lines per tissue. Soft tissue, endometrium, and pancreas have less than 1,000 drug combination tests within less than 3 cell lines per tissue. (c) Number of tested drugs that overlap between tissues. Drugs for each tissue are rarely overlapped with drugs for other tissues except the brain, hematopoietic and lymphoid tissue, bone, and lung. (d) Distribution of monotherapy sensitivity (i.e., relative inhibition) (e) Distribution of combination synergy score (f) Distribution of sensitivity and synergy. Concentrated and dense spikes lie in relative inhibition between 10-50, implying that many experiments have been performed within the relative inhibitions between 10-50. Loose spikes lie in relative inhibition below 0. Here we set the relative inhibition’s binarization threshold at 50 as synergy scores show different distribution after 50.

**Figure 2.** Drug synergy prediction model. (a) Drug encoder. It learns an embedding representation of a drug. Inputs are MACCS fingerprints, canonical SMILES, and target genes. (b) Cell line encoder. It learns an embedding representation of a cell line. Inputs are cancer type and gene expressions. (c) Merging drug encoder and cell line encoder. Sensitivity was an auxiliary output to boost synergy prediction.

**Figure 3.** Transfer learning for understudied tissue. (a) Transfer model parameters from data-rich tissues to data-poor tissue such as bone and prostate (b) Train/test split for data-rich tissues and data-poor tissue. Cell lines from different tissues are disjoint. Some cell lines from different tissues sometimes share drugs.